CESIUM 123

6. POTENTIAL FOR HUMAN EXPOSURE

6.1 OVERVIEW

Stable cesium has been identified in at least 10 of the 1,585 hazardous waste sites that have been proposed for inclusion on the EPA National Priorities List (NPL) (HazDat 2001). It was reported that ¹³⁴Cs has been identified in at least 3 of the 1,585 hazardous waste sites and ¹³⁷Cs has been identified in at least 22 of the 1,585 hazardous waste sites proposed for inclusion on the EPA NPL. However, the number of sites evaluated for cesium is not known. The frequency of these sites within the United States can be seen in Figures 6-1, 6-2, and 6-3. Of these sites, none are located in the Commonwealth of Puerto Rico.

Naturally-occurring cesium and cesium minerals consist of only one stable isotope, ¹³³Cs. Cesium occurs in the earth's crust at low concentrations. Granites contain an average cesium concentration of about 1 ppm and sedimentary rocks contain about 4 ppm (Burt 1993). Higher concentrations are found in lepidolite, carnallite, muscovite, beryl, spodumene, potassium feldspars, leucite, petalite, and related minerals. The most important source of commercial cesium is the mineral pollucite, which usually contains about 5–32% Cs₂O (Burt 1993). The largest deposits of pollucite are located in Manitoba, Canada and account for about two-thirds of the world's known supply. Smaller deposits are located in Zimbabwe, Namibia, Brazil, Scandinavia, Czechoslovakia, and the United States. Continental dust and soil erosion are the main emission sources of naturally occurring cesium present in the environment. Cesium is also released to the environment as a result of human activities. The mining of pollucite ores and the production and use of cesium compounds in the electronic and energy production industries contribute to its direct release to the environment. Cesium has also been detected in the fly ash of hazardous waste incinerators and coal burning power plants (Fernandez et al. 1992; Mumma et al. 1990). Since the production and use of cesium compounds are limited, and since the natural concentration of cesium in the earth's crust is low, ¹³³Cs is not often monitored or detected in the environment.

Of much greater concern is the release of radioactive forms of cesium to the environment, such as ¹³⁷Cs and ¹³⁴Cs. These and other radioactive isotopes were released to the environment as a result of atmospheric testing of nuclear weapons (carried out from 1945 to 1980) and accidents that occurred at nuclear power plants such as the incident at the Chernobyl nuclear power plant in 1986 and the accident at the Windscale nuclear weapons facility in the United Kingdom in 1957. Small amounts of ¹³⁷Cs and ¹³⁴Cs are also released in the airborne and liquid effluents during the normal operation of nuclear power

Figure 6-1. Frequency of NPL Sites with Cesium Contamination



Figure 6-2. Frequency of NPL Sites with Cesium 134 Contamination



Figure 6-3. Frequency of NPL Sites with Cesium 137 Contamination



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plants. These levels are very low in comparison to the amounts released from weapons tests and accidents at nuclear power plants and are not expected to have a major impact upon human health. For the most part, testing of nuclear weapons has been discontinued by most nations for many years now. However, India and Pakistan have recently (May of 1998) conducted limited underground tests of nuclear weapons (UN 1998).

Radioactive material is commonly referred to by its activity rather than its mass. The activity is the number of disintegrations that the material undergoes in a given period of time. The most common units of activity are the curie (Ci) or the becquerel (Bq). One curie is equal to 3.7x10¹⁰ disintegrations per second (dps) and 1 becquerel is equal to 1 dps. For convenience, picocuries (pCi) are often reported for lower activities; 1pCi=1x10⁻¹² Ci (see Section 1.2).

Radioactive cesium is removed from the air by wet and dry deposition and can travel thousands of miles before settling to earth. Wet deposition is considered the most important pathway for the removal of radioactive cesium from the atmosphere. It is a complex process that depends upon meteorological conditions such as temperature, the microphysical structure of the clouds, and the rainfall rate, as well as the physical and chemical properties of the airborne cesium.

Cesium has very low mobility in soil. In general, it has been reported that cesium usually does not migrate below depths of about 40 cm, with the majority of cesium being retained in the upper 20 cm of the soil surface (Korobova et al. 1998; Takenaka et al. 1998). Clay minerals and soils rich in exchangeable potassium adsorb cesium by binding the cations to interlayer positions of the clay particles (Paasikallio 1999). The low hydration energy of cesium cations is primarily responsible for their selective sorption and fixation by clays. These factors can limit the uptake of cesium in grass and plant material. There are exceptional areas, however, where cesium fixation in soil is much less, resulting in greater transport in the soil and uptake in plants. Regions in Venezuela, Brazil, and Russia have been identified where the mobility of cesium is considerably greater than in other soils (LaBrecque and Rosales 1996; WHO 1983). Cesium is also deposited on plants and trees by wet and dry deposition and is absorbed into the flora through its foliage (Sawidis et al. 1990). The deposited cesium can make its way to soil through decomposition of the contaminated foliage.

Since the half-life for some radioactive isotopes of cesium is long (the half-life of ¹³⁷Cs is about 30 years and the half-life of ¹³⁴Cs is about 2 years), the general population is exposed to ¹³⁷Cs and ¹³⁴Cs for long periods of time after it is released from a nuclear accident or weapons test, with the greatest exposure

occurring near the source. Although inhalation and dermal exposure is possible, oral ingestion of contaminated food items is the greatest source of internal exposure for both naturally occurring and radioactive cesium. Cesium is uniformly distributed throughout the whole body similar to potassium, and it does not accumulate in any one particular part of the body like iodine (thyroid) or strontium (bones). For this reason, radioactive cesium only poses significant health risks if a large amount has been ingested. Workers employed in the mining and milling of pollucite ores and the production of cesium compounds are exposed to cesium through oral, dermal, and inhalation routes. Similar routes of exposure to ¹³⁷Cs and ¹³⁴Cs occurs for workers employed in the nuclear industry. External exposure to gamma radiation can also occur for workers employed in the nuclear industry as well as for the general population. The health consequences of external exposure to gamma radiation are not unique to ¹³⁷Cs and ¹³⁴Cs, but are similar for all gamma emitting radionuclides.

6.2 RELEASES TO THE ENVIRONMENT

Throughout this chapter, the units used to express concentration or intake of cesium are the same units reported by the authors. In most cases, values are expressed in mass units when referring to ¹³³Cs, while radioactive cesium isotopes are expressed in units of activity.

According to the Toxic Release Inventory (TRI), in 1999, there were no reportable releases of ¹³³Cs, or its compounds into the environment by commercial sources (TRI99 2001). The TRI data should be used with caution since only certain types of facilities are required to report. Therefore, it is not an exhaustive list. Facilities are required to report to TRI if they have 10 or more full-time employees, or if the facility is classified under Standard Industrial Classification (SIC) codes 20–39, if the facility manufactures or processes over 25,000 pounds of the chemical, or otherwise uses more than 10,000 pounds of the chemical in a calendar year.

In the United States, commercial nuclear power plant operators are required to report any detectable quantities of radioactive materials released to the environment (10 CFR 50.36a). Table 6-1 summarizes releases of ¹³⁷Cs and ¹³⁴Cs to the atmosphere and water for 1993 from pressurized water reactor (PWR) and boiling water reactor (BWR) nuclear power plants. Nearly all of the radioactive material reported as being released in effluents, are from planned releases. Planned releases result from normal plant operation or from anticipated operational occurrences. The latter include unplanned releases of radioactive materials from miscellaneous actions such as equipment failure, operator error, or procedure error; these releases are not of such consequence as to be considered an accident.

Table 6-1. Radiocesium Releases from Nuclear Power Plants for 1993

		Annı	ual total site enviro	onmental releas	es for 1993
			Water		Air
Installation	Location ^a	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci
Boiling water reactors					
Browns Ferry ^b	Decatur, AL	0.033	0.18	3.5x10 ⁻⁴	8.9x10 ⁻⁴
Brunswick ^b	Wilmington, NC	2.9x10 ⁻⁴	1.7x10 ⁻³	No data	7.0x10 ⁻⁴
Clinton	Clinton, IL	No data	No data	No data	No data
Cooper	Omaha, NE	9.3x10 ⁻³	0.052	No data	No data
Dresden ^a	Joliet, IL	1.2x10 ⁻⁶	0.025	No data	1.9x10 ⁻⁴
Duane Arnold	Cedar Rapids, IA	No data	No data	No data	1.4x10 ⁻⁶
Edwin I. Hatch	Baxley, GA	6.3x10 ⁻³	0.044	No data	6.2x10 ⁻⁵
Fermi	Laguna Beach, MI	No data	8.3x10 ⁻⁶	No data	No data
Grand Gulf	Vicksburg, MS	3.5x10 ⁻⁴	6.0x10 ⁻⁴	2.1x10 ⁻⁶	2.3x10 ⁻⁶
Hope Creek	Wilmington, DE	No data	4.8x10 ⁻⁵	No data	No data
Humbolt Bay⁵	Eureka, CA	No data	8.8x10 ⁻³	No data	3.2x10 ⁻⁵
James A. Fitzpatrick	Syracuse, NY	No data	4.1x10 ⁻⁵	No data	6.6x10 ⁻⁶
LaCrosse ^b	LaCrosse, WI	3.2x10 ⁻⁴	0.010	No data	1.0x10 ⁻⁴
LaSalle	Ottawa, IL	No data	No data	No data	No data
Limerick	Philadelphia, PA	2.0x10 ⁻³	6.0x10 ⁻³	No data	No data

Table 6-1. Radiocesium Releases from Nuclear Power Plants for 1993 (continued)

		Annı	ual total site envir	onmental release	es for 1993
			Nater		Air
Installation	Location ^a	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci
Boiling water reactors (cont.)					
Millstone	New London, CT	0.066	0.224	6.9x10 ⁻⁶	8.9x10 ⁻⁵
Monticello	St. Cloud, MN	No data	No data	No data	4.8x10 ⁻⁴
Nine Mile Point	Oswego, NY	No data	No data	No data	1.9x10 ⁻⁵
Oyster Creek	Toms River, NJ	No data	No data	No data	6.5x10 ⁻⁵
Peach Bottom	Lancaster, PA	5.2x10 ⁻⁴	1.4x10 ⁻³	6.1x10 ⁻⁵	3.0x10 ⁻⁴
Perry	Painesville, OH	8.8x10 ⁻⁵	1.6x10 ⁻⁴	No data	No data
Pilgram	Boston, MA	No data	9.8x10 ⁻⁴	2.4x10 ⁻⁵	7.3x10 ⁻⁵
Quad-Cites	Moline, IL	No data	3.6x10 ⁻³	No data	1.8x10 ⁻⁴
River Bend	Baton Rouge, LA	2.6x10 ⁻⁴	2.2x10 ⁻³	No data	No data
Shoreham	Brookhaven, NY	No data	No data	No data	No data
Susquehanna	Berwick, PA	No data	2.9x10 ⁻⁵	No data	No data
Vermont Yankee	Brattleboro, VT	No data	No data	No data	9.9x10 ⁻⁵
WNP-2	Richland, WA	6.3x10 ⁻³	0.019	6.7x10 ⁻⁶	7.2x10 ⁻⁶
Total		0.12	0.58	4.5x10 ⁻⁴	3.3x10 ⁻³

Table 6-1. Radiocesium Releases from Nuclear Power Plants for 1993 (continued)

		Annı	ual total site envir	onmental release	es for 1993
			Nater		Air
nstallation	Location ^a	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci
ressurized water reactors					
Arkansas One	Russellville, AR	0.054	0.11	No data	3.5x10 ⁻⁸
Beaver Valley	Shippingport, PA	1.2x10 ⁻³	2.3x10 ⁻³	No data	No data
Big Rock Point	Charlevoix, MI	9.5x10 ⁻⁴	0.012	1.1x10 ⁻⁵	1.5x10 ⁻⁴
Braidwood	Joliet, IL	6.5x10 ⁻⁴	1.7x10 ⁻³	No data	No data
Byron	Byron, IL	5.6x10 ⁻³	1.5x10 ⁻³	No data	No data
Callaway	Fulton, MO	5.3x10 ⁻⁴	7.8x10 ⁻⁴	No data	No data
Calvert Cliffs	Washington, DC	0.14	0.21	3.4x10 ⁻³	4.2x10 ⁻³
Catawba	Rock Hill, SC	1.6x10 ⁻³	3.9x10 ⁻³	No data	No data
Comanche Peak	Glen Rose, TX	8.7x10 ⁻³	8.7x10 ⁻³	No data	No data
Crystal River	Tampa, FL	2.7x10 ⁻³	9.3x10 ⁻³	No data	3.1x10 ⁻⁶
Davis-Besse	Toledo, OH	1.4x10 ⁻³	6.2x10 ⁻³	No data	No data
Diablo Canyon	San Luis Obispo, CA	0.036	0.064	3.3x10 ⁻⁵	2.4x10 ⁻⁴
Donald C. Cook	St. Joseph, MI	1.3x10 ⁻³	6.3x10 ⁻³	No data	7.3x10 ⁻⁷
Fort Calhoun	Omaha, NE	1.3x10 ⁻³	0.012	No data	1.7x10 ⁻⁶
H.B. Robinson	Hartsville, SC	3.6x10 ⁻⁴	4.1x10 ⁻⁴	5.8x10 ⁻⁶	1.8x10 ⁻⁵
Haddam Neck	Middletown, CT	0.013	0.023	8.0x10 ⁻⁴	8.8x10 ⁻³

Table 6-1. Radiocesium Releases from Nuclear Power Plants for 1993 (continued)

		Annı	ual total site envir	onmental releas	es for 1993
			Water		Air
Installation	Location ^a	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci
Pressurized water reactors (con	nt.)				
Harris	Raleigh, NC	2.9x10 ⁻⁴	3.9x10 ⁻⁴	No data	No data
Indian Point ^b	Peekskill, NY	3.0x10 ⁻³	0.10	No data	7.0x10 ⁻⁴
Joseph M. Farley	Dothan, AL	1.5x10 ⁻³	3.6x10 ⁻³	No data	No data
Kewaunee	Green Bay, WI	No data	2.6x10 ⁻⁶	No data	2.0x10 ⁻⁶
Maine Yankee	Wicassett, ME	1.4x10 ⁻³	8.2x10 ⁻³	No data	4.2x10 ⁻⁵
McGuire	Charlotte, NC	1.6x10 ⁻³	4.6x10 ⁻³	No data	1.3x10 ⁻⁶
North Anna	NW Richmond, VA	6.2x10 ⁻³	9.2x10 ⁻³	3.4x10 ⁻⁶	8.2x10 ⁻⁵
Oconee	Greenville, SC	2.7x10 ⁻³	0.010	No data	3.9x10 ⁻⁴
Palisades	South Haven, MI	2.4x10 ⁻⁴	4.3x10 ⁻³	No data	1.1x10 ⁻⁵
Palo Verde	Phoenix, AZ	No data	No data	2.0x10 ⁻³	1.7x10 ⁻³
Point Beach	Manitowoc, WI	0.019	0.027	6.9x10 ⁻³	6.9x10 ⁻³
Prairie Island	Minneapolis, MN	2.8x10 ⁻³	3.8x10 ⁻³	1.5x10 ⁻⁵	1.8x10 ⁻⁵
R.E. Ginna	Rochester, NY	0.042	0.041	No data	5.2x10 ⁻⁶
Rancho Secob	Sacramento, CA	1.9x10 ⁻⁵	3.6x10 ⁻⁴	No data	No data
Salem	Wilmington, DE	0.81	1.0	No data	7.0x10 ⁻⁷
San Onofre ^b	San Clemente, CA	0.49	0.57	2.5x10 ⁻⁵	4.0x10 ⁻⁵
Seabrook	Portsmouth, NH	No data	3.3x10 ⁻⁵	No data	No data

Table 6-1. Radiocesium Releases from Nuclear Power Plants for 1993 (continued)

		Annı	ual total site envir	onmental release	es for 1993
			Water		Air
Installation	Locationa	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci	¹³⁴ Cs, Ci	¹³⁷ Cs, Ci
Pressurized water reactors (con	nt.)				
Sequoyah	Daisy, TN	0.086	0.14	No data	No data
South Texas	Bay City, TX	3.4x10 ⁻³	5.4x10 ⁻³	No data	No data
St. Lucie	Ft. Pierce, FL	0.055	0.083	9.2x10 ⁻⁶	2.1x10 ⁻⁵
Summer	Columbia, SC	1.6x10 ⁻³	3.3x10 ⁻³	2.2x10 ⁻⁵	2.9x10 ⁻⁵
Surry	Newport News, VA	7.6x10 ⁻⁵	0.011	No data	7.2x10 ⁻⁵
Three Mile Island ^b	Harrisburg, PA	0.026	0.030	1.2x10 ⁻⁷	4.7x10 ⁻⁶
Trojan⁵	Portland, OR	1.0x10 ⁻³	4.0x10 ⁻³	No data	No data
Turkey Point	Florida City, FL	9.4x10 ⁻⁴	5.5x10 ⁻³	No data	9.4x10 ⁻⁷
Vogtle	Augusta, GA	5.6x10 ⁻³	7.3x10 ⁻³	No data	No data
Waterford	New Orleans, LA	0.013	0.016	No data	No data
Wolf Creek	Burlington, KS	0.022	0.024	No data	No data
Yankee Rowe ^b	Greenfield, MA	4.3x10 ⁻⁶	6.0x10 ⁻⁵	No data	1.0x10 ⁻⁷
Zion	Waukegan, IL	0.014	0.029	2.4x10 ⁻⁴	2.9x10 ⁻⁴
Total		1.88	2.85	0.013	0.023

^aPost office state abbreviations used.

^bFacilities that are permanently or indefinitely shut down.

6.2.1 Air

Stable cesium is introduced into the atmosphere by resuspension of soil, accidental release from mining and milling pollucite, and emissions from hazardous waste incinerators or coal burning plants. These emissions are expected to be low since cesium occurs naturally in the earth's crust at low concentrations and only small amounts of pollucite are mined annually. Cesium was detected at concentrations of 10.8 and 6.11 mg/m³ in the effluent of a coal-burning power plant in the western United States (Ondov et al. 1989) and has been identified in the fly ash from municipal incinerators (Mumma et al. 1990, 1991). Fly ash from five municipal waste incinerators in the United States contained cesium at concentrations of 2,100–12,000 ppm (EPA 1990a). Stable cesium has been identified in air at 2 of the 10 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001).

Radioactive isotopes of cesium such as ¹³⁷Cs and ¹³⁴Cs have been released to the atmosphere from atmospheric nuclear weapons testing, accidents at nuclear reactors, and nuclear-powered satellites burning up in the atmosphere upon re-entry. The total amount of ¹³⁷Cs released from weapons testing through 1980 was estimated as 2.6x10⁷ Ci (9.6x10¹⁷ Bq), 76% of which was released in the northern hemisphere and 24% in the southern hemisphere (WHO 1983). On April 26, 1986, a steam buildup caused an explosion and fire at a nuclear power generating plant in Chernobyl, Russia, releasing an estimated $5.4 \times 10^5 \text{ Ci} (2.0 \times 10^{16} \text{ Bg}) \text{ of } ^{134} \text{Cs} \text{ and } 1.1 \times 10^6 \text{ Ci} (4.0 \times 10^{16} \text{ Bg}) \text{ of } ^{137} \text{Cs} \text{ into the atmosphere over Europe}$ (Watson 1987). Long-range transport spread the radionuclides throughout the Northern Hemisphere. No airborne activity from Chernobyl has been reported south of the equator (Eisler 1995). By early May 1986, these radionuclides were readily detectable in environmental samples collected in North America (Huda et al. 1988). More recent estimates have put the total activity of ¹³⁷Cs released from the Chernobyl power plant as 2.3×10^6 Ci $(8.5 \times 10^{16} \text{ Bq})$ and 1.2×10^6 Ci $(4.4 \times 10^{16} \text{ Bq})$ for 134 Cs (Buzulukov and Dobrynin 1993). On January 24, 1978, the Soviet nuclear-powered satellite Cosmos 954 re-entered earth's atmosphere over the Canadian Arctic, releasing an estimated 86 Ci of ¹³⁷Cs (Barrie et al. 1992). In October 1957, an accident at the Windscale nuclear weapons plant at Sellafield in the United Kingdom resulted in the release of 595 Ci of ¹³⁷Cs (ATSDR 1999). Routine activities at nuclear power plants and fuel-reprocessing stations also release ¹³⁷Cs and ¹³⁴Cs to the environment on a regular basis. Radiocesium released in airborne effluents from the normal operation of nuclear power plants is considered low in comparison to releases from atmospheric weapons testing and the major releases following accidents at nuclear power plants. In 1998, it was reported that 1.3x10⁻⁴ Ci of ¹³⁴Cs and 5.1x10⁻³ Ci of ¹³⁷Cs were released to the atmosphere from the Savannah River plutonium processing site in South Carolina (DOE 1998b). In 1993, the Nuclear Regulatory Commission (NRC) estimated that 0.013 Ci of ¹³⁴Cs and

0.023 Ci of ¹³⁷Cs were released in airborne effluents from 30 PWR nuclear power plants operating in the United States (NRC 1993b). It was also estimated that 4.6x10⁻⁴ Ci of ¹³⁴Cs and 3.3x10⁻³ Ci of ¹³⁷Cs were released in airborne effluents from 28 BWR nuclear power plants (NRC 1993b). The total airborne and liquid releases of ¹³⁴Cs and ¹³⁷Cs from the individual nuclear power plants are summarized in Table 6-1.

Radioactive ¹³⁴Cs was not identified in air at the 3 NPL hazardous waste sites where it was detected in some environmental media, but ¹³⁷Cs was identified in air at 5 of the 22 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001).

6.2.2 Water

Cesium can be released to water surfaces during the mining, milling, and production process of pollucite ore. The natural erosion and weathering of rocks will also lead to cesium's introduction into ground and surface water. Stable cesium was identified in groundwater at 4 sites and surface water at 1 of the 10 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001).

The dumping of high and low level radioactive wastes into the Arctic waters by the former Soviet Union has also led to the release of ¹³⁷Cs and ¹³⁴Cs as well as other radioactive nuclides into these waters. In the past, the majority of radioactive cesium released to water surfaces in North America arose from deposition following atmospheric nuclear weapons testing conducted by the United States, primarily during the 1960s (Robbins et al. 1990). Radioactive cesium can be introduced to water from nuclear power plants (accidents and normal operation) and at facilities that produce weapons grade plutonium and uranium. During the period of 1961–1973, it was estimated that about 514 Ci of ¹³⁷Cs was emitted to the Savannah River watershed due to the activities at the Savannah River plutonium processing plant (Olsen et al. 1989). It was further noted that about 18% of this total (92 Ci) drained directly into the Savannah River (Olsen et al. 1989). In 1998, it was reported that 1.0x10⁻⁴ Ci of ¹³⁴Cs and 0.19 Ci of ¹³⁷Cs were released in liquid effluents from the Savannah River plutonium processing site in South Carolina (DOE 1998b). In 1993, the NRC estimated that 1.88 Ci of ¹³⁴Cs and 2.85 Ci of ¹³⁷Cs were released in liquid effluents from 30 PWR nuclear power plants operating in the United States (NRC 1993a). It was also estimated that 0.12 Ci of ¹³⁴Cs and 0.58 Ci of ¹³⁷Cs were released in liquid effluents from 28 BWR nuclear power plants (NRC 1993a). The EPA reported that the total on-site liquid discharges of ¹³⁷Cs from containment ponds at the Nevada Test Site was 0.0017 Ci in 1997 (EPA 1999c). It was estimated that 1,622 Ci of ¹³⁷Cs and 811 Ci of ¹³⁴Cs were released to the cooling pond surrounding the Chernobyl nuclear power plant following the accident in 1986 (UNSCEAR 1996).

Radioactive ¹³⁴Cs was not identified in any water samples at the 3 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001). However, ¹³⁷Cs was identified in groundwater at 4 sites and surface water at 3 of the 22 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001).

6.2.3 Soil

Anthropogenic sources of ¹³³Cs releases to soils include the mining, milling, and processing of pollucite ore. It is also found in the ash of coal burning power plants and municipal waste incinerators. Stable cesium was detected at concentrations of 0.44–2.01 ppm in the bottom ash of municipal solid waste incinerators operating in the United States (Mumma et al. 1990) and at concentrations of 3–23 ppm from a municipal waste incinerator operating in Barcelona, Spain (Fernandez et al. 1992). Stable cesium was identified in soil at 1 site and sediment at 1 of the 10 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001).

Radioactive cesium has been released to soil surfaces by underground nuclear weapons testing, fallout from the accident at the Chernobyl nuclear power plant and fallout from atmospheric weapons testing. About 1,400 underground tests have been performed worldwide, with a total explosive yield of about 90 megatons (ATSDR 1999). Small amounts of ¹³⁷Cs and ¹³⁴Cs are also released to soil from the normal operation of nuclear power plants and the storage of spent fuel rods. Not including the 30-km exclusionary zone, an area of approximately 2.4×10^4 km² near the Chernobyl nuclear power plant was contaminated with ¹³⁷Cs at a deposition density >5.4x10⁻⁵ Ci/m² following the accident in 1986 (UNSCEAR 1996). Within the exclusionary zone the contamination density may have been 2 orders of magnitude greater in limited areas (UNSCEAR 1996). The mean deposition density of ¹³⁷Cs and ¹³⁴Cs in four different soils in Devoke, United Kingdom for May 1986 were reported as 3.7x10⁻⁷-5.4x10⁻⁷ Ci/m² and 1.0x10⁻⁷–1.8x10⁻⁷ Ci/m², respectively (Hilton et al. 1993). The concentrations of ¹³⁷Cs in eight sediment cores of the Danube River, Austria were about 540 pCi/kg in April 1985, but increased to approximately 27,000-81,000 pCi/kg in October 1986, following the accident at the Chernobyl nuclear power plant (Rank et al. 1990). The deposition of ¹³⁷Cs attributed to the accident at the Chernobyl nuclear power plant in sediment at five different sites in Lake Constance, Germany ranged from 2.7x10⁻⁷ to 2.1x10⁻⁶ Ci/m², while the concentration attributed to fallout from nuclear weapons testing since 1963 ranged from 1.4x10⁻⁷ to 5.4x10⁻⁷ Ci/m² (Richter et al. 1993). It was estimated that 2,973 Ci of ¹³⁷Cs and 1,622 Ci of ¹³⁴Cs were released to the sediments in the cooling pond surrounding the Chernobyl nuclear power plant following the accident in 1986 (UNSCEAR 1996). The deposition density of ¹³⁷Cs in

123 soil cores collected at the Idaho National Engineering and Environmental Laboratory (INEEL), a site for stored transuranic waste in the United States, ranged from 1.6x10⁻⁸ to 3.4x10⁻⁷ Ci/m² (DOE 1998a). The deposition density of ¹³⁷Cs in soils from Idaho, Montana, and Wyoming ranged from 3.0x10⁻⁹ to 1.1x10⁻⁷ Ci/m², and it was assumed that its origin was fallout from the Nevada Test Site (DOE 1998a). The mean deposition density of ¹³⁷Cs in the top layer (0–8 cm) of soils near the Chernobyl nuclear power plant in 1988 was 8.6x10⁻⁵ Ci/m² and the mean deposition density of ¹³⁴Cs was 1.9x10⁻⁵ Ci/m² (Mikhaylovskaya et al. 1993).

It was reported that ¹³⁴Cs was identified in soil at 1 site, but was not detected in any sediment samples at the 3 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001). It was also reported that ¹³⁷Cs was identified in soil at 13 sites and sediment at 5 of the 22 NPL hazardous waste sites where it was detected in some environmental media (HazDat 2001).

6.3 ENVIRONMENTAL FATE

6.3.1 Transport and Partitioning

The transport and partitioning of particulate matter in the atmosphere is largely dependent upon the physical properties of the matter such as size and density as well as the meteorological conditions such as temperature, the microphysical structure of the clouds, and rainfall rate. The particle size of ¹³⁷Cs released to the atmosphere following the Chernobyl nuclear accident was in the range of 0.1–10 μm, (Hirose et al. 1993). Particles <5 μm in diameter usually have low deposition velocities and are transported long distances before being removed from the atmosphere. Atmospheric cesium is deposited on land and water via wet and dry deposition and the deposited cesium may be re-suspended to the atmosphere by disturbances that occur on the ground such as vehicular traffic and construction activity. The wet deposition velocity of ¹³⁷Cs at Tsukauba, Japan from May 5 to May 30, 1986 ranged from 0.0026 to 0.110 m/second, and the largest value recorded was during a period of heavy rainfall (Hirose et al. 1993). The mean deposition velocity (wet and dry) of ¹³⁷Cs measured in Prague, Czechoslovakia was reported as 0.08 m/second from 1989–1992 and the mean flux rate was 1,108 pCi/m²-year (Rybacek et al. 1994).

Since cesium does not volatilize from water, transport of cesium from water to the atmosphere is not considered likely, except by windblown sea sprays. Most of the cesium released to water will adsorb to suspended solids in the water column and ultimately be deposited in the sediment core. Cesium can also

bioconcentrate and has been shown to bioaccumulate in both terrestrial and aquatic food chains. Mean bioconcentration factors (BCF) for ¹³⁷Cs of 146, 124, and 63 were reported for fish, brown macroalgae, and molluscs, respectively (Fisher et al. 1999). Mean BCF values of 92, 58, 39, and 150 were reported for ¹³⁷Cs in cod, haddock, plaice, and whiting, respectively (Steele 1990). In a study of aquatic organisms inhabiting the Ottawa River, a 4-fold increase of ¹³⁷Cs levels was observed with each trophic level (Rowan et al. 1998). The levels of ¹³⁷Cs in lake trout from Great Slave Lake, Canada were consistently higher than levels found in food sources and a biomagnification factor of 1.9 was calculated for lake trout, relative to their food sources. The biomagnification factor was 3.5 for large mature trout populating the lake (Rowan et al. 1998). It was shown that the bioconcentration and bioaccumulation of ¹³⁷Cs by aquatic organisms is significantly reduced in waters with a large humic content and high levels of potassium cations (Penttila et al. 1993). Because of the high potassium concentration in oceans, the transfer of ¹³⁷Cs and ¹³⁴Cs to fish is much greater in freshwater and the activity of freshwater fish may be 100 times that of ocean fish, given the same cesium concentration in the water (WHO 1983).

In soil surfaces, cesium has low mobility in comparison to other metals and usually does not migrate below a depth of 40 cm. The majority portion of cesium is retained in the upper 20 cm of the soil surface (Korobova et al. 1998; Ruse and Peart 2000; Takenaka et al. 1998). Vertical migration patterns of ¹³⁷Cs in four agricultural soils from southern Chile indicated that approximately 90% of the applied cesium was retained in the top 40 cm of soil, and that in one soil, essentially 100% was bound in the upper 10 cm (Schuller et al. 1997). Migration rates of radiocesium were derived from the depth distribution profiles and were in the range of 0.11 to 0.29 cm/year (Schuller et al. 1997). The vertical migration patterns of ⁹⁰Sr and ¹³⁷Cs produced from the atomic bomb exploded in Nagasaki, Japan were studied over a 40-year period (Mahara 1993). Over this period, 95% of the cesium remained in the top 10 cm of the soil surface and no cesium was detected below a depth of 40 cm. In contrast, only 70% of 90 Sr was located within a depth of 10 cm and a small percentage was detectable below a depth of 200 cm. The in situ vertical migration rate of ⁹⁰Sr was calculated as 0.42 cm/year and the migration rate of ¹³⁷Cs was 0.10 cm/year (Mahara 1993). Soil adsorption coefficients (K_d) of five radionuclides (⁵⁴Mn, ⁶⁰Co, ⁶⁵Zn, ⁸⁵Sr, and ¹³⁷Cs) were measured for 36 agricultural soils collected in Japan. It was determined that ¹³⁷Cs had the largest median K_d of all five radionuclides, and that a positive correlation was observed between the adsorption coefficient and exchangeable potassium content in the soil (Yasuda et al. 1995). No correlations were observed for other soil properties such as pH, water content, cation exchange capacity, and exchangeable calcium. Other studies have reported that clay and zeolite minerals strongly bind cesium cations and can therefore reduce the bioavailability of cesium and the uptake in plants by irreversibly binding cesium in interlayer positions of the clay particles (Paasikallio 1999). Experiments conducted by growing plants in a peat soil showed that the introduction of zeolites into the soil-plant system decreased the uptake of ¹³⁴Cs

in plants by a factor of 8 (Shenber and Johanson 1992). The low hydration energy of cesium cations is primarily responsible for their selective sorption and fixation by clays and zeolites (Hakem et al. 1997). Soils rich in organic matter adsorb cesium, but the cesium adsorbed in the organic fraction is readily exchangeable and highly available for plant uptake (Sanchez et al. 1999). Regions in Venezuela, Brazil, and Russia have been identified where a great deal of rain is encountered, the soil is peaty or podzolic (a type of forest soil characterized by high leachability), and the mobility of cesium is considerably greater than in other soils (LaBrecque and Rosales 1996; WHO 1983).

The plant/soil concentration ratio (activity/kg of plant/activity/kg of soil) of ¹³⁷Cs for field crops in southern Finland ranged from 0.01 to 0.26. In northern Finland, this ratio ranged from 0.01 to 2.29, with the lowest values occurring in clay and silt soils (Paasikallio et al. 1994). The plant/soil concentration ratios for a series of vegetables and grains decreased in the following order: lettuce, cabbage>carrot, potato>cereals, onion; for fruits, the order was: blackcurrant>strawberry>apple (Paasikallio et al. 1994). The mean plant/soil concentration ratios of ¹³⁷Cs for trees at the Hanford Waste Site in the United States were 0.03 (roots), 0.06 (cores), and 0.02 (leaf/twig) (Landeen and Mitchell 1986).

6.3.2 Transformation and Degradation

6.3.2.1 Air

When pure cesium metal is exposed to air, an explosion-like oxidation occurs, forming a mixture of cesium oxides (Cs₂O, Cs₂O₂, and Cs₂O₃). Cesium compounds released to the atmosphere will eventually settle to earth by wet and dry deposition. Radioactive forms of cesium such as ¹³⁷Cs and ¹³⁴Cs are continuously transformed to stable isotopes of barium or xenon by the natural process of radioactive decay. The pathways and mechanisms of these reactions have been described in Chapter 4.

6.3.2.2 Water

When pure cesium metal is released to water, a vigorous reaction occurs yielding cesium hydroxide (CsOH), the strongest base known, and hydrogen gas, which may ignite spontaneously. In general, cesium compounds are very water soluble, and exist primarily as the Cs⁺ cation. Under normal environmental conditions, Cs⁺ cations are neither degraded nor transformed, but may adsorb to suspended solids and sediment in the water column, forming insoluble complexes.

6.3.2.3 Sediment and Soil

Cesium salts and most cesium compounds are generally very water soluble, with the exception of cesium alkyl and aryl compounds, which have low water solubility. Cesium cations have a low hydration energy and can react with clay minerals, zeolites, or soils with a high percentage of exchangeable potassium, forming insoluble, immobile complexes.

6.4 LEVELS MONITORED OR ESTIMATED IN THE ENVIRONMENT

Reliable evaluation of the potential for human exposure to cesium depends in part on the reliability of supporting analytical data from environmental samples and biological specimens. Radioactive cesium is measured in units of activity, not mass. A great deal of monitoring data for radioactive cesium in environmental samples prior to and shortly following the accident at the Chernobyl nuclear power plant on April 26, 1986 have been included.

6.4.1 Air

Data reporting the background levels of ¹³³Cs in the atmosphere are limited. Since the quantity of cesium mined and milled is small, it is expected that background concentrations in the atmosphere will be low. The concentration of ¹³³Cs measured in the South Pole during 1974–1975 was reported in the range of 0.072 to 0.14 pg/m³, with a mean of 0.10 pg/m³ (Maenhaut et al. 1979). The maximum airborne concentrations of ¹³³Cs at the Tera Nova Bay Station in the Arctic were reported as 20–90 pg/m³ during the 1989–1990 Italian expedition and 10–60 pg/m³ for the 1990–1991 expedition (Chiavarini et al. 1994). The average concentration of ¹³³Cs in precipitation collected in Lennox, Massachusetts during the summer of 1984 was reported as 0.0075 µg/L, with a range of 0.0050 to 0.024 µg/L (Dasch and Wolff 1989).

Radioactive ¹³⁴Cs and ¹³⁷Cs have been detected at various concentrations (activities) in the atmosphere following the accident at the Chernobyl nuclear power plant on April 26, 1986. The greatest concentrations were observed in locations throughout Russia and Europe, but detectable levels were measured globally, including in North America. The concentrations of ¹³⁴Cs in the air above the destroyed reactor were 1,756.8 pCi/m³ on May 8 and 54.0 pCi/m³ on May 18 (Buzulukov and Dobrynin 1993). The concentrations of ¹³⁷Cs in the air above the destroyed reactor were 8,918.9 pCi/m³ on May 8 and 135.1 pCi/m³ on May 18 (Buzulukov and Dobrynin 1993). Atmospheric concentrations of ¹³⁴Cs in Belgrade, Yugoslavia were 5.4x10⁻⁴ (May 1, 1986), 160.5 (May 2, 1986), 145 (May 3, 1986), and

25.4 (May 4, 1986) pCi/m³, while atmospheric concentrations of ¹³⁷Cs were 0.001 (May 1, 1986), 324 (May 2, 1986), 276 (May 3, 1986), and 49 (May 4, 1986) pCi/m³ (Ajdacic and Martic 1990). These concentrations decreased significantly over time as advection, wet deposition, and dry deposition lowered the atmospheric concentrations. The average monthly atmospheric concentration of ¹³⁷Cs in Belgrade that were measured during 1991–1996 ranged from 1.3x10⁻⁴ to 2.0x10⁻³ pCi/m³, with a pronounced maximum during the winter months (Todorovic et al. 1999). Monitoring data from Prague, Czechoslovakia indicated that there was a gradual decrease in the atmospheric concentration of ¹³⁷Cs from May 1986 to February 1992. Maximum concentrations were measured immediately following the accident at the Chernobyl nuclear power plant (approximately 0.05 pCi/m³) and gradually decreased over a 6-year period (Rybacek et al. 1994). The atmospheric concentration of ¹³⁴Cs decreased more rapidly, presumably due to the shorter half-life of ¹³⁴Cs. The concentration of ¹³⁷Cs in the atmosphere of Thessaloniki, Greece ranged from 8.1x10⁻⁴ to 0.044 pCi/m³ from July 1987 to December 1988, and the concentration of ¹³⁷Cs in rainfall ranged from 0.27 to 34 pCi/L from November 1986 to February 1989 (Papastefanou et al. 1989). During a heavy rainfall event occurring on May 6, 1986, ¹³⁷Cs was detected at a concentration of 46,000 pCi/L (Papastefanou et al. 1989). The concentration of ¹³⁷Cs in Tsukuba, Japan during May 1986 ranged from about 0.054 to 1.6 pCi/m³ (Hirose et al. 1993). The average atmospheric concentrations of ¹³⁴Cs and ¹³⁷Cs in eastern Canada were reported as 0.024 and 0.046 pCi/m³, respectively, during May 10–24, 1986 (Huda et al. 1988). The maximum atmospheric concentration of ¹³⁷Cs measured in New York City in May 1986 was reported as 0.26 pCi/m³ (Feely et al. 1988). The average concentrations of ¹³⁷Cs and ¹³⁴Cs in Barrow, Alaska were reported as 0.027 and 0.019 pCi/m³, respectively, for the month of May 1986 (DOE 1986). In 1975, the maximum concentration of ¹³⁷Cs in the atmosphere, in Poland was 1.89 pCi/m³ (Glowiak et al. 1977b).

6.4.2 Water

The background concentration of 133 Cs in fresh water lakes and rivers is ordinarily in the range of 0.01 to 1.2 µg/L, and the concentration in seawater is about 0.5 µg/L (WHO 1983). Stable cesium was detected in streams that feed into the Tamagawa River, Japan at concentrations of $9x10^{-4}$ to 0.093 µg/L (Tanizaki et al. 1992). Studies from five distinct geochemical areas of the semi-arid endorheic region of the Famatina Range (La Rioja, Argentina) have shown some areas contain high levels of 133 Cs in natural waters and sediment (Fernandez-Turiel et al. 1995). The cesium concentration in fresh water systems of this region ranged from 0.58 to 3.69 µg/L (Fernandez-Turiel et al. 1995). The concentrations of 19 trace metals were studied in drinking water and groundwater supplies in southern Nigeria. Stable cesium was detected in groundwater and drinking water at mean concentrations of 0.61 µg/L (range 0.09–3.72 µg/L)

and $0.35 \mu g/L$ (range 0.05– $4.32 \mu g/L$), respectively (Asubiojo et al. 1997). It was further noted that the mean concentration of cesium in drinking water was lower than the mean concentration of any of the other trace elements analyzed.

High and low level radioactive wastes have been dumped by the former Soviet Union into remote Arctic waters, leading to the release of radioactive cesium into the Kara and Barents Seas. The level of ¹³⁷Cs in surface water of the Barents Sea and Kara Sea was 0.14 and 0.16 pCi/L, respectively, and it was also detected in deep water of the Barents Sea at a concentration of 0.15 pCi/L (Fisher et al. 1999). The concentration of ¹³⁷Cs in the Black Sea was in the range of 2.7 to 8.1 pCi/L for the period of 1991–1996, with the exception of the spring of 1992, when concentrations as high as 43 pCi/L were observed (Strezov et al. 1999). From 1988 to 1991, the mean concentrations of ¹³⁷Cs and ¹³⁴Cs along the Spanish coast of the Mediterranean Sea were 0.13 and 0.0072 pCi/L, respectively (Molero et al. 1999). Due to its shorter half-life, ¹³⁴Cs was detected in all 14 samples collected in 1988 and 1989, but only in 3 samples collected in 1990 and 1991, suggesting that the ¹³⁴Cs levels observed in surface Mediterranean waters during this period were due exclusively to Chernobyl-related deposition. The ¹³⁷Cs concentration incorporated into the Mediterranean Sea near the Spanish coast from the post-Chernobyl fallout was about 0.032 pCi/L, which was approximately a 33% increase over previous levels (Molero et al. 1999). Maximum ¹³⁷Cs and ¹³⁴Cs levels in the immediate vicinity of nuclear power plants located on the southern Catalan shore of the Mediterranean were 0.57 and 0.059 pCi/L (Molero et al. 1999). Concentrations of ¹³⁷Cs in lakes and streams in Devoke, United Kingdom decreased exponentially from a maximum concentration of about 8.1 pCi/L on May 6, 1986 to about 0.027 pCi/L 1,200 days later (Hilton et al. 1993). The mean concentration of ¹³⁷Cs in six lakes located in central Finland ranged from 111 pCi/L in 1987 to 8.1 pCi/L in 1989 (Penttila et al. 1993).

The concentration of ¹³⁷Cs and ¹³⁴Cs in groundwater at 18 U.S. Department of Energy (DOE) facilities was reported in the range of 2.7x10⁻³ to 1.83x10³ pCi/L (DOE 1992). The concentration of ¹³⁷Cs measured in groundwater wells at Carlsbad, New Mexico (the site of Project GNOME) ranged from 99 to 6.8x10⁵ pCi/L in 1997 (EPA 1999c). The concentration of ¹³⁷Cs in groundwater at the Chernobyl nuclear power plant was in the range of 40.5–1,100 pCi/L in 1988 and 29.7–129.7 pCi/L in 1989 (Prister et al. 1990). The mean concentration of ¹³⁷Cs in drinking water in Poland in 1974 was reported as 0.2 pCi/L (Glowiak et al. 1977b). These concentrations in water may be compared to the federal radiation safety standards. For continuous ingestion over a lifetime, the maximum concentrations of ¹³⁴Cs and ¹³⁷Cs in drinking water are limited to 900 and 1,000 pCi/L respectively. It should be noted however, that these

limits assume no other intake of radioactivity. If other radioisotopes are ingested, then the intake limits for each must be adjusted proportionately (NRC 1999a).

6.4.3 Sediment and Soil

Stable cesium occurs naturally in the earth's crust at low concentrations. Granites contain an average cesium concentration of about 1 ppm and sedimentary rocks contain about 4 ppm (Burt 1993). Others have estimated cesium concentration of granites as high as 5 ppm (WHO 1983). Stable cesium was detected in dust samples from roadside and pedestrian traffic in Nagpur, India at concentrations of 1.53–3.63 μg/g, with the largest value obtained in the vicinity of a metals factory (Chutke et al. 1995). It was reported that ¹³³Cs was detected at concentrations ranging from 0.9 to 2.2 μg/g in alluvial sediments in the Sava River, Croatia (Vertacnik et al. 1997). The concentration range of ¹³³Cs in river sediment from five distinct geochemical areas of the semi-arid endorheic region of the Famatina Range (La Rioja, Argentina) was 2.28–6.20 μg/g (Fernandez-Turiel et al. 1995).

The concentration of ¹³⁷Cs in soils of Thessaloniki, Greece ranged from 1,440 to 35,324 pCi/kg (average 8,154 pCi/kg) and the concentration of ¹³⁴Cs ranged from about 270 to 5,676 pCi/kg during the period of August 1986 to February 1989, with most of the fallout attributed to the accident at the Chernobyl nuclear power plant (Papastefanou et al. 1989). The concentration of ¹³⁷Cs in 10 uncultivated fields from southern England ranged from 0 to 946 pCi/kg, with the highest levels contained in the top 10 cm of the soil surface (Owens et al. 1996). The concentration of ¹³⁷Cs in five cultivated fields ranged from 0 to 540 pCi/kg, and the concentrations were well distributed from the surface to the plough layer (Owens et al. 1996). The concentration of ¹³⁷Cs in three soils in Hong Kong receiving a large amount of rainfall ranged from 32 to 201 pCi/kg (Ruse and Peart 2000). The concentration of ¹³⁷Cs in sediment from the Ribble Estuary, England near the British Nuclear Fuels Laboratory ranged from 270 to 1,351 pCi/kg (Brown et al. 1999). The average concentration of ¹³⁷Cs in sediment from Lake Ontario was reported as 8,108 pCi/kg, and suspended solids from the Niagara River contained 324 pCi/kg (Platford and Joshi 1989). The average concentrations of ¹³⁷Cs in uncultivated soils in northern Poland were reported as 616–4,170 pCi/kg from 1988–1991 (Pietrzak-Flis et al. 1994). The mean concentration of ¹³⁷Cs in surface soil samples from the Los Alamos nuclear laboratory test site during the period of 1974–1996 was reported as 611 pCi/kg (Fresquez et al. 1998). Concentrations of ¹³⁷Cs around the perimeter of the site and background concentrations off the site were reported as 589 and 419 pCi/kg, respectively. The concentration of ¹³⁷Cs and ¹³⁴Cs in soils and sediments at 18 U.S. DOE facilities was reported to range from 20 to 4.69x10⁷ pCi/kg (DOE 1992). The concentration of ¹³⁷Cs in sediment from

the Savannah River ranged from 5 to 100 pCi/kg in 1986, while the concentration in suspended solids and particulate matter ranged from 240 to 4.324x10⁶ pCi/kg (Olsen et al. 1989). The mean concentration of ¹³⁷Cs in soil at the Hanford Site in the United States was 4,540 pCi/kg (Landeen and Mitchell 1986). The mean concentration of ¹³⁷Cs in soils taken from two high-elevation sites in northern Colorado ranged from 4,054 to 7,027 pCi/kg (Ulsh et al. 2000).

6.4.4 Other Environmental Media

Data regarding the concentrations of ¹³⁷Cs, ¹³⁴Cs, and ¹³³Cs in various animals are shown in Table 6-2. Concentrations are dependent on the location, date, and level of exposure. For example, the concentrations of ¹³⁷Cs in bullhead catfish inhabiting an abandoned nuclear reactor reservoir at the Savannah River site in South Carolina were as high as 1.54x10⁵ pCi/kg (McCreedy et al. 1997), but concentrations in various freshwater species of fish in the Ottawa River were in the range of 54 to 351 pCi/kg (Rowan et al. 1998). After the accident at the Chernobyl nuclear power plant, the average concentrations of ¹³⁷Cs in perch, pike, and roach obtained from 52 freshwater lakes in Finland were 55,811, 66,297, and 15,270 pCi/kg, respectively, in 1988. By contrast, the mean concentrations in 1992 had fallen to 14,324, 18,567, and 5,892 pCi/kg, respectively (Sarkka et al. 1996). The concentration of total ¹³⁷Cs and ¹³⁴Cs measured in body tissue from sheep in Ireland from 1989 to 1991 ranged from about 2,700 to 10,811 pCi/kg, with highest levels observed during November of 1989 (McGee et al. 1993).

The concentrations of ¹³³Cs in a series of plants from Britain were studied over a 1-year period. Concentrations of 50–300 ng/g were measured, with the highest levels observed during the summer and fall months. The median concentration of ¹³³Cs in poplar leaves collected in Bulgaria was reported as 75 ng/g, while concentrations in land plants ranged from 30 to 440 ng/g (Djingova et al. 1995). In contrast to the radioactive isotopes of cesium, concentrations of ¹³³Cs depend primarily on the root uptake from soil and not on atmospheric deposition. Lichens and mosses have been shown to trap and retain ¹³⁷Cs and ¹³⁴Cs more so than vascular plants, due to their relatively large surface area. Lichens and mosses from northern Greece contained ¹³⁷Cs levels of 6.6x10⁴–5.1x10⁵ pCi/kg during the period of 1989–1991 (Papastefanou et al. 1992) and moss samples from Finland collected in 1988–1989 contained 4.3x10⁴–9.7x10⁵ pCi/kg (Penttila et al. 1993). The mean concentration of ¹³⁷Cs in three species of lichens collected in August 1986 from Megalopolis, Greece were 2.6x10⁴–3.3x10⁴ pCi/kg, while the mean concentrations for the same three species of lichens collected in October 1996 had fallen to 3,324–7,892 pCi/kg (Riga-Karandinos and Karandinos 1998). Mushrooms, lichens, and mosses obtained near Manitoba, Canada in August 1986 contained ¹³⁷Cs at mean concentrations of 6.4x10⁵, 8.6x10⁴, and

Table 6-2. Concentration of ¹³³Cs, ¹³⁴Cs, and ¹³⁷Cs in Animals

Species	Location and date	¹³³ Cs (µg/g)	134Cs (Bq/kg)	¹³⁷ Cs (Bq/kg)	Source
Royal albatross (N=3)	Indian Ocean (1994)	0.009-0.025 (0.014 mean); liver	No data	No data	Kim et al. 1998
Black-footed albatross (N=18)	North Pacific (1985)	0.007-0.049 (0.022 mean); liver	No data	No data	Kim et al. 1998
Black-browed albatross (N=9)	Indian Ocean (1994)	0.013–0.042 (0.022 mean); liver	No data	No data	Kim et al. 1998
White-capped albatross (N=3)	Indian Ocean (1994)	0.011–0.039 (0.029 mean); liver	No data	No data	Kim et al. 1998
Yellow-nosed albatross (N=4)	Indian Ocean (1994)	0.0029–0.079 (0.022 mean); liver	No data	No data	Kim et al. 1998
Grey-headed albatross (N=10)	Indian Ocean (1994)	0.011–0.031 (0.016 mean); liver	No data	No data	Kim et al. 1998
Northern giant petrel (N=6)	Indian Ocean (1994)	0.005-0.034 (0.015 mean); liver	No data	No data	Kim et al. 1998
Northern fulmar (N=17)	North Pacific (1985)	0.008–0.036 (0.016 mean); liver	No data	No data	Kim et al. 1998
Bluefin tuna (N=14)	Newfoundland (1990)	0.08–0.24 (0.13 mean); muscle	No data	No data	Hellou et al. 1992a
Cod (N=12)	Newfoundland (1990)	0.16–0.24 (0.19 mean); muscle	No data	No data	Hellou et al. 1992b
Cod (N=10)	Newfoundland (1990)	0.14–0.36 (0.23 mean); muscle	No data	No data	Hellou et al. 1992b
Pilot whale (N=9)	North Atlantic (1987–1996)	0–0.010 (0.006 mean); liver	No data	No data	Becker et al. 1997
White-sided dolphin (N=4)	North Atlantic (1987–1996)	0.027–0.042 (0.032 mean); liver	No data	No data	Becker et al. 1997

Table 6-2. Concentration of ¹³³Cs, ¹³⁴Cs, and ¹³⁷Cs in Animals (*continued*)

Species	Location and date	¹³³ Cs (μg/g)	¹³⁴ Cs (Bq/kg)	¹³⁷ Cs (Bq/kg)	Source
Beluga whale (N=15)	Arctic (1987–1996)	0.021–0.046 (0.031 mean); liver	No data	No data	Becker et al. 1997
Ringed seal (N=13)	Arctic (1987–1996)	0.0045–0.048 (0,020 mean); liver	No data	No data	Becker et al. 1997
Woodcock (N=24)	Ireland (1986)	No data	3.9-206.4; muscle	6.2-565.5; muscle	Pearce 1995
Duck (N=5)	Ireland (1986)	No data	2.2-14.3; muscle	6.4-18.0; muscle	Pearce 1995
Snipe (N=5)	Ireland (1986–1987)	No data	1.0-5.4; muscle	3.6-16.9; muscle	Pearce 1995
Reindeer (N=8)	Northern Sweden (1986–1987)	No data	No data	900 (mean); muscle	Ahman and Ahman 1994
Deer (N=11)	Los Alamos (1991–1998)	No data	No data	2,516 (mean); muscle	Fresquez et al. 1999a
Deer (N=11)	Los Alamos (1991–1998)	No data	No data	888 (mean); bone	Fresquez et al. 1999a
Caribou (N=18)	Saskatchewan (1995)	No data	No data	58 (mean); bone	Thomas and Gates 1999
Caribou (N=18)	Saskatchewan (1995)	No data	No data	228 (mean); liver	Thomas and Gates 1999
Caribou (N=18)	Saskatchewan (1995)	No data	No data	367 (mean); muscle	Thomas and Gates 1999
Caribou (N=18)	Saskatchewan (1995)	No data	No data	553 (mean); kidney	Thomas and Gates 1999
Caribou (N=36)	Alaska (1987)	No data	No data	26–232; neck	Allaye-Chan et al. 1990
Caribou (N=36)	Alaska (1987)	No data	No data	28.4-201.1; shoulder	Allaye-Chan et al. 1990
Caribou (N=36)	Alaska (1987)	No data	No data	30.2-166.5; back	Allaye-Chan et al. 1990

8.4x10⁴ pCi/kg, respectively (Mihok et al. 1989). Since caribou and reindeer consume large amounts of this vegetation during the winter months, high levels of ¹³⁷Cs and ¹³⁴Cs have been detected in these animals.

Stable cesium has been detected infrequently in food products at low concentrations. The average concentration of 133 Cs in 110 onion samples collected in Denmark was 0.21 µg/kg, with a range of not detected to 0.98 µg/kg (Bibak et al. 1998). By comparison, other elements such as calcium and potassium had mean concentrations of $2x10^5$ and $1.6x10^6$ µg/kg, respectively. The concentration range of 133 Cs in wheat flour samples collected in Pakistan was 6.7–11.2 ppb, but 133 Cs was not detected from wheat flour samples from America (Ahman et al. 1994).

Levels of ¹³⁷Cs were below detection limits for all foods analyzed for in the U.S. Food and Drug Administration (FDA) Total Diet Study in 1991–1996 with the exception of honey (Capar and Cunningham 2000). The concentration of ¹³⁷Cs in honey from the 1995 Market Basket Survey was 6.7 Bg/kg (181.1 pCi/kg), which is almost 200 times lower than the regulatory level of ¹³⁷Cs in foods. The average concentrations of total ¹³⁷Cs and ¹³⁴Cs in milk powder, infant milk powder, infant cereal, meat, lentil, wheat, and macaroni samples from Saudi Arabia were 514, 351, 486, 162, 270, 676, and 351 pCi/kg, respectively (Abdul-Majid et al. 1992). For the month of June 1986, the average concentrations of total ¹³⁷Cs and ¹³⁴Cs in milk, green vegetables, fruit, lamb, and beef were reported as 3,243, 2,703, 2,703, 8,108, and 1,622 pCi/kg in high deposition areas of the United Kingdom (Cumbria, north Wales, Scotland, northern Ireland, and the Isle of Man) (Mondon and Walters 1990). It was also estimated that the concentration of total ¹³⁷Cs and ¹³⁴Cs was <676 pCi/kg in each of these food sources in areas of low deposition during this time frame. The maximum concentration of ¹³⁷Cs in pasteurized milk from 65 cities in the United States was 14 pCi/L in May 1989 (EPA 1989). The concentration of ¹³⁷Cs in fresh milk from Chester, New York and pasteurized milk samples from New York City in May 1986 ranged from about 5.4 to 18.9 pCi/L (Feely et al. 1988). Using radiological surveys from 1978 and 1985–1986, the concentration of ¹³⁷Cs in 44 adult food groups from the Rongelap Island and Rongelap Atoll was in the range of 0.52–13,000 pCi/kg (Robinson and Phillips 1989).

6.5 GENERAL POPULATION AND OCCUPATIONAL EXPOSURE

General population exposure to cesium occurs by ingestion of food and drinking water, by inhalation of ambient air, and dermal contact with cesium in soil. Since 133Cs is found in low concentrations in the environment, the exposure to the general population is expected to be low. The National Health And Nutritional Examination Survey (NHANES), conducted by the Centers for Disease Control (CDC), reported that the geometric mean concentration of ¹³³Cs in the urine of 1,006 U.S. residents was 4.7 μg/L in 1999 (CDC 2001). Since the sample size of the 1999 survey was small and was conducted in only 12 locations across the United States, the CDC will combine 1999 and year 2000 data and publish an updated report in the future. Occupational exposure to ¹³³Cs occurs primarily through inhalation and dermal contact at workplaces where pollucite is mined or cesium compounds are manufactured or used. The National Occupational Exposure Survey (NOES) conducted by the National Institute for Occupational Safety and Health (NIOSH) from 1981 to 1983 has estimated that 16,461 workers (4,276 of these were female) were potentially exposed to cesium and cesium compounds in the United States (NOES 1989). The NOES database does not contain information on the frequency, level, or duration of the exposure of workers to any of the chemicals listed therein. They are surveys that only provide estimates of workers potentially exposed to the chemicals. The median concentration of ¹³³Cs in the lungs of metal workers in northern Sweden was 0.016 µg/g and was lower than the median concentration of 0.021 µg/g for a control group that was not occupationally exposed (Hewitt 1988). The range of ¹³³Cs concentrations in lung tissue of coal miners from the United Kingdom was 0.07–0.91 µg/g (Hewitt 1988).

Exposure to radioactive cesium is more important from a health perspective than exposure to stable cesium. The primary source of radioactive cesium in the environment is due to fallout from past atmospheric nuclear weapons tests and the Chernobyl accident. Additional contributions from the normal operation of nuclear power plants and other nuclear facilities are small by comparison. Current exposure of the general U.S. population to ¹³⁴Cs and ¹³⁷Cs is expected to be low since atmospheric testing of nuclear weapons has been discontinued for many years and Chernobyl-related fallout was low in the United States. As discussed in Appendix A, the average annual effective dose of ionizing radiation (including ¹³⁴Cs and ¹³⁷Cs) to the U.S. population from anthropogenic sources are negligible in comparison to natural sources, especially radon and its decay products.

The average daily intake (AVDI) of ¹³⁷Cs and ¹³⁴Cs was estimated for adult males residing in the Ukraine in 1994, based upon total diet samples. The mean intake of ¹³⁷Cs was estimated as 109 pCi/day and the mean intake of ¹³⁴Cs was estimated as 8.1 pCi/day (Shiraishi et al. 1997). Based on dietary patterns and

the concentration of radiocesium in food sources, the total dietary intakes of ¹³⁴Cs and ¹³⁷Cs for typical adults residing in Croatia for the month of May 1986 were estimated as 2.8x10⁴ and 5.9x10⁴ pCi, respectively (Lokobauer et al. 1988). The mean monthly levels of ¹³⁷Cs in human muscle tissue in Graz, Austria were reported as 1,519 (July 1986 to June 1987), 1,049 (July 1987 to June 1988), 340.5 (July 1988 to June 1989), and 202.7 pCi/kg (July 1989 to June 1990), with a maximum value of 9,584 pCi/kg observed in an individual during September 1986 (Rabitsch et al. 1991). The monthly averages for ¹³⁴Cs were about half of those reported for ¹³⁷Cs. By comparison, the maximum concentration of ¹³⁷Cs in muscle tissue from Harwell, England in 1959 was reported as 224 pCi/kg, the mean concentration in muscle tissue from Massachusetts during January 1961 to June 1962 was 100 pCi/kg, and the mean concentration in human muscle tissue obtained from Japan during April to December 1963 was reported as 119 pCi/kg (Rabitsch et al. 1991). The mean concentrations of ¹³⁷Cs in the urinary excretion of Italians in northern Italy were 7.3 and 6.2 pCi/day in 1995 and 1996, respectively (Ropolo and Cesana 1997). These values were about two orders of magnitude less than values reported for measurements taken in 1987. The mean concentration of ¹³⁷Cs in brain, heart, liver, gonads, muscle, bone, and teeth were 0.440, 1.860, 0.490, 2.440, 0.017, 0.106, and 0.23 pCi/g, respectively, for adult cadavers over 34 years of age in Poland during 1975 (Glowiak et al. 1977a). The mean body burdens of ¹³⁷Cs for adults in Helsinki, Viitasaari and Ammans, Finland from 1987 to 1994 are given in Table 6-3 (Rahola and Suomela 1998).

Persons employed at nuclear power facilities and waste disposal sites are potentially exposed to higher levels of ¹³⁷Cs and ¹³⁴Cs than the general population. The NOES conducted by NIOSH from 1981 to 1983 estimated that 13,148 workers (1,294 of these were female) were potentially exposed to ¹³⁴Cs and ¹³⁷Cs in the United States (NOES 1989).

6.6 EXPOSURES OF CHILDREN

This section focuses on exposures from conception to maturity at 18 years in humans. Differences from adults in susceptibility to hazardous substances are discussed in 3.7 Children's Susceptibility.

Children are not small adults. A child's exposure may differ from an adult's exposure in many ways. Children drink more fluids, eat more food, breathe more air per kilogram of body weight, and have a larger skin surface in proportion to their body volume. A child's diet often differs from that of adults.

Table 6-3. The Mean Body Burdens of ¹³⁷Cs for Adults in Finland from 1987–1994^a

Location	Date	Concentration (pCi/kg)
Helsinki, Finland	1987	649
	1988	568
	1989	405
	1990	297
	1991	246
	1992	214
	1993	195
	1994	181
Viitasaari, Finland	1987	3,514
	1988	1,946
	1989	1,595
	1990	1,081
	1991	838
	1992	676
	1993	649
	1994	514
Ammans, Finland	1987	2,892
	1988	3,108
	1989	2,243
	1990	2,486
	1991	1,568
	1992	1,405
	1993	865
	1994	811

^aRahola and Suomela 1998

^{***}DRAFT FOR PUBLIC COMMENT***

The developing human's source of nutrition changes with age: from placental nourishment to breast milk or formula to the diet of older children who eat more of certain types of foods than adults. A child's behavior and lifestyle also influence exposure. Children crawl on the floor, put things in their mouths, sometimes eat inappropriate things (such as dirt or paint chips), and spend more time outdoors. Children also are closer to the ground, and they do not use the judgment of adults to avoid hazards (NRC 1993a).

As for adults in the general population, exposures of children to cesium occur from normal ingestion of food and drinking water, inhaling air, and dermal contact with cesium in soil. No information on cesium levels in amniotic fluid, meconium, cord blood, or neonatal blood was available.

Radioactive cesium was detected in several pasteurized milk and breast milk samples worldwide and since children tend to consume large amounts of milk, this represents an important source of childhood exposure. The maximum concentration of ¹³⁷Cs in pasteurized milk from 65 cities in the United States was 14 pCi/L in May 1989 (EPA 1989). Concentrations of ¹³⁷Cs in human milk samples from several U.S. cities from 1956 to 1961 were <20 pCi/L (Eaman 1986). The overall ¹³⁷Cs concentration in milk taken 7 days postpartum from 37 mothers in two Italian hospitals were 5.8–115 pCi/L (Eaman 1986). In a study of females from northern Sweden, ¹³⁷Cs was detected in breast milk from 8 out of 12 mothers at concentrations of 7.3–178.4 pCi/kg (Johansson et al. 1998). The infants of these mothers who were breast-fed had whole-body levels of ¹³⁷Cs in the range of 45.9–675.7 pCi/kg (Johansson et al. 1998). Based on dietary patterns and the concentration of radiocesium in food sources, the total dietary intakes of ¹³⁴Cs and ¹³⁷Cs for children (10-years-old) residing in Croatia for May–June 1986 were estimated as 43,000 and 190,000 pCi, respectively (Lokobauer et al. 1988). For infants (1 year of age), it was estimated that the total intakes of ¹³⁴Cs and ¹³⁷Cs were 46,000 and 170,000 pCi, respectively (Lokobauer et al. 1988). The total intakes of ¹³⁴Cs and ¹³⁷Cs for adults during this same time period were estimated as 40,000 and 84,000 pCi, respectively (Lokobauer et al. 1988). The higher intakes of ¹³⁴Cs and ¹³⁷Cs for infants and children were traced to a much greater consumption of contaminated milk.

The tendency of young children to ingest soil, either intentionally through pica or unintentionally through hand-to-mouth activity, is well documented. These behavioral traits can result in ingestion of cesium present in soil and dust. Soil ingestion may be a potentially important exposure pathway in areas that have historically had a great deal of ¹³⁴Cs and ¹³⁷Cs deposited onto soil surfaces from the accident at the Chernobyl nuclear power plant or fallout from weapons testing. Playing in contaminated soil could also lead to dermal and external exposure. Ingested cesium is adsorbed strongly to soils and may not be in bioavailable form. A study in which 102 healthy volunteers were fed ¹³⁴Cs-contaminated soil pellets,

only about 1% of the original amount was absorbed and on average, about 60% of the intake was excreted within 48 hours (LeRoy et al. 1966).

It is unlikely that children whose parents are employed at nuclear power generating plants and facilities that store or handle radioactive waste are exposed to ¹³⁴Cs and ¹³⁷Cs from parents' clothing or items removed from the workplace because exit monitors exist at nuclear power plants, and the extensive use of outerware that remains in the plant to prevent these types of incidents. It is also unlikely that children are exposed to ¹³³Cs from parents' clothing or items that have been removed from the workplace if the parents are employed in the mining, milling, or processing of pollucite ore. Other home exposures are unlikely since household items or products used in crafts, hobbies, or cottage industries do not contain significant amounts of cesium.

6.7 POPULATIONS WITH POTENTIALLY HIGH EXPOSURES

Few populations are likely to be exposed to high levels of ¹³³Cs. Persons residing in the vicinity of pollucite mines and workers employed in the mining, milling, and production of cesium may be exposed to higher levels than the general population.

Human exposure to ¹³⁴Cs and ¹³⁷Cs can be external, due to exposure from a radioactive cloud and contaminated environmental media after deposition, or internal, via inhalation and ingestion of contaminated food or drinking water. In both cases, populations residing near the source are exposed to potentially higher doses than populations far removed from the source. Humans residing near areas where nuclear weapons testing was previously conducted may have been exposed to higher doses of radiation from ¹³⁴Cs and ¹³⁷Cs, both internally and externally, than the general population. Populations residing in southern Utah and Nevada were exposed to radioactive cesium (and many other radionuclides) due to testing conducted at the Nevada Test Site (NTS). A total of 100 surface or near-surface tests with a total explosive yield of about 1 megaton were performed at the Nevada test site between 1951 and 1962. The dust from these tests also drifted over the continental United States, producing varying degrees of exposure for remote populations depending upon the meteorological conditions. For example, the greatest non-local fallout from one of the tests was received in New York state in 1953, some 2,000 miles away from the source, due to wet deposition (ATSDR 1999). About 500 underground tests were also performed at the NTS, but underground testing rarely leads to significant off-site contamination (ATSDR 1999). The EPA Office of Radiation and Indoor Air conducts off-site environmental monitoring around former U.S. nuclear test areas. The 1997 report concluded that the current exposure to populations

around the NTS from radionuclides, including ¹³⁷Cs and ¹³⁴Cs, was negligible in comparison to background levels (EPA 1999c).

Populations residing in the vicinity of nuclear power plants may also be exposed to higher levels of ¹³⁴Cs and ¹³⁷Cs than the general population due to airborne and liquid effluents from these plants. Persons employed in these facilities are also likely to be exposed to greater levels than the general population, particularly those employees who must handle radioactive waste material. However, despite the potential for exposure, increased body burdens of radioactive cesium have not been observed among the population of workers in nuclear facilities.

Human populations that received a large amount of fallout from the Chernobyl nuclear accident are also potentially exposed to high levels of radiocesium. These areas were primarily located in the Ukraine and northern Europe that received a great deal of rainfall in the weeks following the accident. Not including the 30-km exclusionary zone, an area of approximately 2.4x10⁴ km² near the plant was contaminated with 137 Cs at a deposition density >5.4x10⁻⁵ Ci/m² (UNSCEAR 1996). Within the exclusionary zone, the contamination density may have been a factor of about 100 greater in limited areas (UNSCEAR 1996). The Bryansk-Belarus region, about 200 km northeast of the reactor, had deposition densities as high as 1.3x10⁻⁴ Ci/m² and the Kaluga-Tula-Orel location, approximately 500 km northeast of the reactor had deposition densities of about 1.6x10⁻⁵ Ci/m² (ATSDR 1999). While cesium is not considerably taken up by the roots of vascular plants, the deposition of radioactive debris on flora with large surface areas such as lichens or moss is significant (see Section 6.4.4). As a result, animals that feed on this vegetation such as caribou or reindeer may ingest large quantities of radiocesium. Nordic or Eskimo populations which use these animals as an important source for food are exposed to larger quantities of ¹³⁷Cs and ¹³⁴Cs than the general population (Allaye-Chan et al. 1990; WHO 1983). The average concentration of ¹³⁷Cs in the skeletal muscle of a herd of caribou from Alaska was in the range of 76–133 Bg/kg in 1987 (Allaye-Chan et al. 1990). According to the National Council on Radiation Protection and Measurements, the maximum nonoccupational radiocesium intake is 300,666 Bq/year (8.1x10⁶ pCi/year) for adults (NCRP 1977). At the maximum average skeletal muscle concentration (133 Bg/kg), an annual consumption of over 2,260 kg would be required to reach this limit (assuming no other intake sources). Using the mean ¹³⁷Cs level of 900 Bg/kg in the muscle of reindeer obtained from northern Sweden in 1986–1987 (Ahman and Ahman 1994), over 330 kg of contaminated meat would have to be consumed to reach the maximum intake level. Populations residing in the Marshall Islands were exposed to higher levels of ¹³⁷Cs than the general population due to nuclear weapons testing conducted by the United States from 1946 to 1958. Using radiological surveys from 1978 and 1985–1986, the AVDI of ¹³⁷Cs due to food ingestion was

estimated for seven age groups residing in the Rongelap Atoll (Robinson and Phillips 1989). The AVDI in pCi/day were as follows: 0–3 months of age, 424; 4–8 months of age, 556; 9 months to 1.4 years of age 773; 1.5–3 years of age, 517; 4–11 years of age, 594; 12–17 years of age, 761; over 18 years of age, 1,085 (Robinson and Phillips 1989). This corresponds to an annual intake of 3.96x10⁵ pCi/year, for adults over 18 years of age.

6.8 ADEQUACY OF THE DATABASE

Section 104(i)(5) of CERCLA, as amended, directs the Administrator of ATSDR (in consultation with the Administrator of EPA and agencies and programs of the Public Health Service) to assess whether adequate information on the health effects of cesium is available. Where adequate information is not available, ATSDR, in conjunction with the National Toxicology Program (NTP), is required to assure the initiation of a program of research designed to determine the health effects (and techniques for developing methods to determine such health effects) of cesium.

The following categories of possible data needs have been identified by a joint team of scientists from ATSDR, NTP, and EPA. They are defined as substance-specific informational needs that if met would reduce the uncertainties of human health assessment. This definition should not be interpreted to mean that all data needs discussed in this section must be filled. In the future, the identified data needs will be evaluated and prioritized, and a substance-specific research agenda will be proposed.

6.8.1 Identification of Data Needs

Physical and Chemical Properties. As shown in Table 4-2, the relevant physical and chemical properties of cesium and its compounds are known (Burt 1993; Lide 1998). In general, cesium compounds are very water soluble and primarily exist in ionic form in aqueous environments. Cesium adsorbs strongly to soils and is not very mobile (Korobova et al. 1998; Paasikallio 1999; Ruse and Peart 2000; Takenaka et al. 1998). The radioactive decay modes of the two most important cesium isotopes, ¹³⁴Cs and ¹³⁷Cs, are also well understood (ICRP 1983) and no physical or chemical data needs are required to permit the prediction of the environmental fate of cesium.

Production, Import/Export, Use, Release, and Disposal. Knowledge of a chemical's production volume is important because it may indicate possible environmental contamination and human exposure. If a chemical's production volume is high, there is an increased probability of general population exposure via consumer products and environmental sources such as air, drinking water, and food. Data concerning production volumes, import, and use of commercially significant cesium compounds are not available. No information was found for cesium in the TRI. According to the Emergency Planning and Community Right-to-Know Act of 1986, 42 U.S.C. Section 11023, industries are required to submit chemical release and off-site transfer information to the EPA. This database will be updated yearly and should provide a list of industrial production facilities and emissions. The United States is 100% import reliant for ¹³³Cs. There are no salient statistics such as production volume, consumption, or import/export volumes of cesium. Although there is no information regarding the countries shipping cesium or cesium compounds to the United States, it is believed that Canada is the major source of cesium (USGS 1999). Data regarding the U.S. production (if any) of cesium and its compounds as well as import/export statistics would be useful. The amount of ¹³⁴Cs and ¹³⁷Cs released in airborne and liquid effluents from nuclear power plants in the United States is known (NRC 1993b).

Environmental Fate. Information is available to permit assessment of the environmental fate and transport of cesium in air (Hirose et al. 1993; Rybacek et al. 1994), water (Burt 1993; WHO 1983), and sediment and soil (Korobova et al. 1998; Paasikallio 1999; Ruse and Peart 2000; Takenaka et al. 1998). Cesium compounds are water soluble, and only one oxidation state (+1) is observed under environmental conditions (Burt 1993). Cesium released to the atmosphere may be carried long distances before being deposited onto soil and water surfaces by wet and dry deposition (Hirose et al. 1993). Most of the cesium released to water will adsorb to suspended solids in the water column and ultimately be deposited in the sediment core. In soil surfaces, cesium is strongly adsorbed in the upper layers and generally has very low mobility (Korobova et al. 1998; Paasikallio 1999; Ruse and Peart 2000; Takenaka et al. 1998). This is particularly true for soils with a high potassium content or soils rich in clay. The radioactive decay modes of the two most important cesium isotopes, ¹³⁴Cs and ¹³⁷Cs, have been described in Chapter 4 (ICRP 1983).

Bioavailability from Environmental Media. The bioavailability of cesium in environmental media is well understood and no data needs are required at this time. For the most part, cesium is adsorbed strongly in the surface of most soils and is not readily bioavailable (Paasikallio 1999). In a study in which 102 healthy volunteers were fed ¹³⁴Cs-contaminated soil pellets, only about 1% of the original intake was absorbed, and on average, about 60% of the original amount was excreted within 48 hours

(LeRoy et al. 1966). In peaty or podzolic soils with a low clay content, cesium is reversibly adsorbed to the organic fraction and is expected to be in bioavailable form (Sanchez et al. 1999). Cesium uptake in vascular plants has been demonstrated (Djingova et al. 1995; Willey and Martin 1995). Increasing the clay or potassium content results in lower uptake by plants (Shenber and Johanson 1992). It has also been shown that fish residing in waters with high concentrations of humic material and potassium, such as oceans, have lower whole-body cesium concentrations than fish in freshwater where the concentration of dissolved potassium is lower given the same cesium concentration in the water (WHO 1983).

Food Chain Bioaccumulation. Cesium bioaccumulates in both aquatic and terrestrial food chains (Rowan et al. 1998; WHO 1983). Mean BCFs for ¹³⁷Cs of 146, 124, and 63 were reported for fish, brown macroalgae, and molluses, respectively (Fisher et al. 1999). Cesium accumulation in aquatic organisms occurs from both food sources and cesium dissolved in the water column or adsorbed to suspended solids and sediments. The lichen-caribou-human food chain is an important route of human exposure and has been well studied (Allaye-Chan et al. 1990; WHO 1983). No additional data are required to assess the potential for human exposure to cesium through food chain bioaccumulation.

Exposure Levels in Environmental Media. Stable cesium and radioactive cesium have been detected in air (Ajdacic and Martic 1990; Chiavarini et al. 1994; Dasch and Wolff 1989; Rybacek et al. 1994; Todorovic et al. 1999), water (Asubiojo et al. 1997; DOE 1992; Fisher et al. 1999; Prister et al. 1990; Strezov et al. 1999; WHO 1983), soil/sediment (Burt 1993; DOE 1992, 1998a; Ruse and Peart 2000; WHO 1983), and food (Ahman et al. 1994; Bibak et al. 1998; Mondon and Walters 1990). Due to the large surface area of lichens and moss, they can collect a great deal of atmospheric nuclear fallout and often have high concentrations of ¹³⁴Cs and ¹³⁷Cs (Papastefanou et al. 1992; Penttila et al. 1993). Grazing animals such as reindeer and caribou that feed on large quantities of lichens or moss may potentially ingest large amounts of radioactive cesium and this may be transferred to humans who consume these animals as a meat source (Allaye-Chan et al. 1990; WHO 1983). Continued monitoring data on ¹³⁴Cs and ¹³⁷Cs in environmental media are needed to extend knowledge of human exposure to these radionuclides.

Exposure Levels in Humans. Monitoring data exist for the levels of ¹³⁴Cs and ¹³⁷Cs in human populations (Glowiak et al. 1977b; Lokobauer et al. 1988; Rabitsch et al. 1991; Rahola and Suomela 1998; Ropolo and Cesana 1997). Most of the current data is for areas of eastern Europe and Russia. Limited data are available regarding the levels of ¹³³Cs in humans (Hewitt 1988). Stable cesium was detected in the urine of U.S. residents at a geometric mean concentration of 4.2 μg/L (CDC 2001). More information regarding the background concentration of ¹³³Cs in human populations would be useful, but

given the ubiquitous distribution of cesium at low levels in the environment, background levels are unlikely to approach levels that would reflect cesium toxicity. Since the levels of radioactivity change over time, continued monitoring data on the levels of ¹³⁴Cs and ¹³⁷Cs in humans are needed in order to interpret health effects that may occur at current exposure levels.

Exposures of Children. Children, like adults, are not expected to be over exposed to ¹³³Cs. Existing data show that children are exposed to ¹³⁴Cs and ¹³⁷Cs after major releases. These exposures are primarily related to dietary intake and the intake of contaminated milk (Lokobauer et al. 1988). Dietary consumption patterns suggest that the weight-adjusted intake of radioactive cesium for children may be greater than for adults following a nuclear release (Lokobauer et al. 1988). In a study of females from northern Sweden, ¹³⁷Cs was detected in breast milk from 8 out of 12 mothers at concentrations of 7.3–178.4 pCi/kg (Johansson et al. 1998). The infants of these mothers who were being breast-fed had whole-body levels of ¹³⁷Cs in the range of 45.9–675.7 pCi/kg (Johansson et al. 1998). Additional data regarding the transfer of ¹³⁴Cs and ¹³⁷Cs from breast milk and contaminated pasteurized milk samples to children, as well as body burden studies on children, would be useful in assessing the potential risk that these radionuclides would pose following a major release.

Child health data needs relating to susceptibility are discussed in Section 3.12.2, Identification of Data Needs: Children's Susceptibility.

Exposure Registries. Cesium is currently one of the chemicals for which a subregistry has been established in the National Exposure Registry. The information that is amassed in the National Exposure Registry facilitates the epidemiological research needed to assess adverse health outcomes that may be related to exposure to cesium.

6.8.2 Ongoing Studies

The database of federal research programs in progress (FEDRIP 2000) indicates several current projects that may fill some existing data gaps and add to the current database of knowledge. Studies are in progress that seek to identify methods for more effective removal of radioactive cesium from contaminated soils and wastes. Phytotech Incorporated is conducting a study regarding the mobilization and removal of strontium and cesium from soil by chemical treatment and phytoremediation. Rufus L. Chaney from the Beltsville Agricultural Research Center in Beltsville, Maryland is researching the feasibility of developing plants, soil, and plant management practices that can cost-effectively

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phytoextract radionuclides and heavy metals from contaminated soils. Dr. Heit from the U.S. DOE is conducting measurements regarding the fallout of nuclear debris from both atmospheric weapons tests and accidental atmospheric releases to determine the mechanisms of transport and deposition and to verify and correct fallout models. Work is being performed by Dr. Kinkead at the Los Alamos National Laboratory regarding the separation of cesium and strontium from high level radioactive waste. Dr. Leon Kochian from the Agricultural Research Center in Ithaca, New York is investigating the bioaccumulation of ¹³⁷Cs in plants grown in contaminated soils.